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1 **Persistent and polarised global actin flow is essential for**
2 **directionality during cell migration**

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18 **Abstract**

19 Cell migration is hypothesised to involve a cycle of behaviours beginning with
20 leading edge extension. However, recent evidence suggests that the leading edge
21 may be dispensable for migration, raising the question of what actually controls cell
22 directionality. Here we exploit the embryonic migration of *Drosophila* macrophages to
23 bridge the different temporal scales of the behaviours controlling motility. This
24 reveals that edge fluctuations during random motility are impersistent and weakly
25 correlated with motion. In contrast, flow of the actin network behind the leading edge
26 is highly persistent. Quantification of actin flow structure during migration reveals a
27 stable organisation and asymmetry in the cell-wide flowfield that strongly correlates
28 with cell directionality. This organisation is regulated by a gradient of actin network
29 compression and destruction, which is controlled by myosin contraction and cofilin-
30 mediated disassembly. It is this stable actin-flow polarity, which integrates rapid
31 fluctuations of the leading edge that controls inherent cellular persistence.

Introduction

Cell migration is hypothesised to involve a stepwise cycle of behaviours starting with protrusion of the leading edge¹. At the same time the cell must maintain polarity, which is hypothesised to be controlled by a combination of reaction-diffusion modules and membrane tension to maintain an asymmetry in these behaviours²⁻⁵. The integration of these stages leads to coherent motion whereby cells have an inherent persistence in speed and direction⁶.

Since the first postulation of the migratory cycle, we now understand many of its molecular components. Protrusion of the leading edge is driven by Arp2/3-mediated actin polymerisation^{7, 8}. Pushing of the actin filaments against the leading edge, along with myosin-II contraction, subsequently induces a retrograde motion of the crosslinked actin-network⁹⁻¹¹. When integrins are engaged, these points of friction with the extracellular matrix lead to a slowing of the retrograde flow allowing for transmission of the forces of the flowing actin network into extracellular traction stresses^{12, 13}. Finally, asymmetric release of integrin adhesions at the rear allows for net cell translocation¹⁴.

This model of migration is predicated on the idea that it is a linear stepwise process starting at the leading edge^{1, 15}. Therefore, with regards to the control of cell directionality, a major focus has been on the actin polymerisation machinery at the front¹⁶. However, evidence has arisen to bring this leading edge centric view into question. Loss of leading edge lamellipodia does not grossly inhibit chemotaxis^{17, 18} and the presence of actin protrusions, rather than being essential for cell translocation, may actually destabilise migration and enhance exploratory behaviour^{19, 20}. Additionally, recent data revealed that cell shape is a predictor of migratory dynamics,^{21, 22} suggesting that global cellular processes are also important

58 in controlling motility. However, the idea that cell movement may not be directly
59 controlled by extensions at the cell edge is still controversial²³ as the stepwise view
60 of migration has remained nearly unchanged for decades²⁴.

61 What is currently lacking is a holistic understanding of how the hypothesised
62 steps of motility are integrated in space and time to give a cell its inherent
63 persistence and directionality. The complexity in bridging the stages of migration is
64 partly due to the different time and lengthscales of these behaviours²⁵. For instance,
65 it is difficult to understand how rapid fluctuation of the leading edge, which oscillates
66 on the order of seconds²⁶⁻²⁸, controls overall cell persistence that decays on the
67 order of minutes to hours⁶. To resolve such questions requires migration to be
68 imaged at sufficiently high spatiotemporal resolution for long time periods to correlate
69 edge fluctuation, actin dynamics, and overall cell motion. However, correlating
70 behaviours on such different timescales is both experimentally and quantitatively
71 challenging.

72 In previous work, we exploited the embryonic migration of *Drosophila*
73 macrophages (hemocytes), which are highly amenable to live imaging during their
74 developmental dispersal²⁹, to develop tools to image cell-wide actin flows during
75 their migration *in vivo*³⁰. Here we use this system to quantify, and correlate in time,
76 the various behaviours of motility during both random and directed migration. This
77 reveals that edge fluctuation is a weak predictor of directionality during random
78 motility, with a persistence that is less than the overall persistence of the cell. In
79 contrast, the retrograde flow of the actin network behind the leading edge, which has
80 recently been revealed to couple cell speed and persistence³¹, is highly organised
81 and stable in time. Through the development of approaches to quantify global actin
82 flow organisation we reveal that cell migration involves network-wide coordination

83 and an intrinsic asymmetry in the flowfield that highly correlates with cell
84 directionality. This asymmetry is controlled by a stable gradient of actin network
85 compression and destruction towards the rear of the lamellae, which is driven by
86 myosin contraction and cofilin-mediated disassembly. It is this
87 destruction/contraction gradient that leads to a stable cell-wide polarity within the
88 flowing actin network, which likely integrates the rapid fluctuations of the leading
89 edge to control overall cell persistence.

90

91

92 **Results**

93 ***Leading edge fluctuations are a weak predictor of cell directionality***

94 Hemocytes can be automatically and precisely tracked during their embryonic
95 migration using the nucleus as a fiducial marker, which is more accurate than
96 tracking cell centroid at high temporal resolution (Supplementary Video 1). We first
97 used the natural variation in the fluctuation of hemocyte contours (Extended Data
98 Fig. 1A) to highlight a relationship between edge activity and cell motion at high
99 temporal resolution (5 s/frame). However, morphodynamic analysis revealed that
100 edge fluctuations are largely disconnected from cell speed (Fig. 1A-F). We next
101 quantified the positions of edge extensions during random migration with respect to
102 the instantaneous direction of motion to determine how protrusions are correlated
103 with cell directionality. Normalising the position vectors of edge extensions or
104 retractions to hemocyte directionality revealed that edge fluctuations are weakly
105 correlated with motion (Fig. 1G-J; Extended Data Fig. 1B; Supplementary Video 2).
106 These data are surprising considering the high persistence of hemocytes
107 (directionality ratio 0.7 ± 0.1 SEM, this work)³². Interestingly, taking into account both

direction and speed of extensions (resultant velocity) revealed that all extensions around the cell collectively showed stronger correlation to motion than the maximum extension alone (Extended Data Fig. 1C); this suggests that minor extensions that individually are not correlated to motion are integrated to provide directional information. Finally, comparing the persistence of cell trajectory and the maximum extension revealed that the leading edge was less persistent than overall cell motion (Fig. 1K). These data highlight that randomly patrolling hemocytes spend significant effort generating extensions independent of motion in a mode of motility that has been termed 'inefficient'^{27, 33}, suggesting that other behaviours must be involved to provide their high migratory persistence.

Actin flow is globally organised in migrating cells

Recent mathematical modelling has suggested that actin flow may help establish the inherent persistence of migrating cells³¹. To examine cell-wide actin flows in hemocytes we performed Particle Image Velocimetry (PIV) of LifeAct-GFP expressing cells. Global PIV analysis suggested an overall organisation to the actin flow with vectors showing a high degree of alignment and a gradient of high to low flow speed from the leading edge to the cell body (Fig. 2A; Supplementary Video 3). As LifeAct-GFP binds actin indirectly, we wanted to confirm that PIV was actually highlighting internal motion of the network. We therefore labelled directly with Actin-GFP, and photobleached spots in the network, which allowed us to examine network transit. Photobleached spots at the leading edge moved in a retrograde fashion toward the cell body and mimicked the flowfield of LifeAct-GFP expressing cells, suggesting PIV analysis was indeed tracking actin motion (Supplementary Video 4). In order to understand how actin flow was structured we first calculated its

divergence, which highlights sources and sinks in the network. While there was little positive divergence (Extended Data Fig. 2A), the network showed large regions of high negative divergence at the rear of the lamella immediately adjacent to the hemocyte cell body (Fig. 2B; Extended Data Fig. 2B; Supplementary Video 3), which correlated with zones of actin fibre deformation (Extended Data Fig. 2C). In contrast to the rapid fluctuation of the leading edge, these negatively divergent regions were persistent on the order of 30-60 seconds (Extended Data Fig. 2B).

In order to examine the global organisation of actin flow we seeded streamlines at each point along the edge of migrating hemocytes. The evolution of streamlines during migration showed an overall organisation in the actin flowfield with many streamlines ending within a region of the lamella anterior to the cell body in the direction of cell travel (Fig. 2C; Supplementary Video 3). We calculated the strength of streamline confluence, which revealed a predominant streamline endpoint that was asymmetrically distributed within the cell (Fig. 2D; Supplementary Video 3). Interestingly, this streamline endpoint was highly negatively divergent and tended to represent a region of low actin flow speed (Fig. 2E; Extended Data Fig. 2D; Supplementary Video 3), showing that it represents a large sink within the actin flowfield. Additionally, the number of sinks appeared to correlate with the number of hemocyte lamellae suggesting that they may contribute to maintaining cell polarity (Extended Data Fig. 2E; Supplementary Video 5).

The primary streamline sink appeared to represent some transition in the actin flow as its location was strongly correlated with sharp transition from retrograde to anterograde actin motion (Fig. 2F-H). We hypothesise that these transition regions are analogous to the retrograde/anterograde transitions observed in migrating cells *in vitro*^{18, 34-36} and the transition of actin network gripping to slipping. When we

calculated the actin flow speed in the retrograde region, this revealed that the actin flow was indeed slower anterior to the primary sink (Extended Data Fig. 2F), suggesting that this is where extracellular friction is highest. While this is consistent with the actin-clutch hypothesis, we observed no relationship between actin flow speed and cell speed as has been hypothesised should occur³¹, suggesting that this linear relationship may not be valid on shorter timescales (Extended Data Fig. 2G).

In order to examine whether actin flow organisation can be observed in other migrating cells we examined fish keratocytes, growth cones^{28, 37}, and retinal pigmented epithelial cells (RPE1) (Extended Data Fig. 3A, Supplementary Videos 6-8). All cell types showed similar global organisation of actin flow with streamlines converging at a large network sink deep within the cell, suggesting that this is a conserved feature of migrating cells.

During random migration, the polarity of global actin flow is highly correlated with hemocyte directionality

We next examined how actin flow organisation correlated with leading edge dynamics and cell directionality. In order to correlate leading edge extension and flow polarity with cell motion, we calculated vectors from the nucleus to defined points within the cell and correlated the direction of these vectors with the cell's direction of motion. This revealed that the primary streamline sink and the retrograde/anterograde transition region were more correlated with cell motion than edge extension/retraction (Fig. 3A; Extended Data Fig. 4A; Supplementary Video 9). Furthermore, the persistence of the primary sink was higher than the persistence of the leading edge (Fig. 3B). These data suggest that the coordinated flow of actin

may be integrating leading edge activity to provide an inherent persistence to randomly migrating hemocytes.

We next tested if there was a temporal hierarchy of these various migratory behaviours. Interestingly, even at our rapid temporal resolution of 5 s/frame we observed a maximum correlation at 0-lag, highlighting that edge extension, actin flow polarity, and cell directionality are precisely correlated in time (Fig. 3C). This is despite the fact that these different phenomena are separated by relatively large distances within the cell (Extended Data Fig. 4B).

We next examined the correlation of leading edge activity and the streamline sink with cell directionality in RPE1 cells. Similar to hemocytes, edge speed in RPE1 cells was uncorrelated with cell speed (Extended Data Fig. 3B). Furthermore, both the direction of the maximum extension and the primary sink were correlated with cell motion (Extended Data Fig. 3C,D; Supplementary Video 10). Additionally, the direction of the velocities of the maximum extension, primary sink, and cell motion showed a maximum correlation at 0-lag, highlighting that, similar to hemocytes, these behaviours were strongly coordinated (Extended Data Fig. 3E). However, in contrast to hemocytes the maximum extension and primary sink were equivalently correlated with cell motion (Extended Data Fig. 3D) suggesting that RPE1 cells are more efficient with regards to the production of edge protrusions.

During directed migration, leading edge persistence controls hemocyte directionality

While the organisation of global actin flow appeared more important than edge fluctuations in defining the persistence of randomly patrolling hemocytes, we wondered whether the same would be true during directed migration. Hemocytes

can be rapidly induced to migrate to epithelial wounds through hydrogen peroxide release³⁸. When migration behaviours were examined during a wound response (Extended Data Fig. 4C; Supplementary Videos 11,12) there was an increase in the correlation of the maximum extension and primary sink with cell motion (Fig. 3D; Extended Data Fig. 4D-F). Furthermore, there was an increase in leading edge persistence, which unlike randomly moving cells, closely matched the persistence of other migratory behaviours (Fig. 3E,F; Extended Data Fig. 4G). Interestingly, cells undergoing directed and random migration showed equivalent distributions of extensions around the cell perimeter when normalised to the direction of motion (Fig. 3G). However, the resultant edge velocity of these extensions was more correlated to motion in directionally migrating cells (Fig. 3H), suggesting that the wound cue may be increasing the speed of edge extensions in the direction of the wound site. These changes in leading edge activity were also correlated with an increase in the persistence of cells migrating to the wound (Extended Data Fig. 4H). These data suggest that the leading edge in hemocytes is more critical for driving migration during chemotactic responses, which is similar to what has been reported for mammalian dendritic cells²⁰.

Nonequilibrium fluid-mechanical model spontaneously breaks symmetry, resulting in highly stable actin flows and persistent motion

To gain insight into the connections between actin flows, sinks, and persistent cell motion, we built a minimal one-dimensional fluid-mechanical model with active processes based on only four coupled partial differential equations (see Supplementary modelling details and related models for more information^{39, 40}). This minimal model leads to an emergent actin flow profile with a gradient of myosin

intensity and a sink at the rear (Fig. 3I). The induction of stable flow was relatively insensitive to changes in actin polymerisation (as long as a threshold was reached; Fig. 3J) or depolymerisation (see Supplementary modelling details) but sensitive to changes in myosin levels, suggesting the flow was remarkably robust (Fig. 3J). Additionally, it was relatively difficult to reorient the flow by perturbation of cortical actin density. When we simulated a single pulse of actin at a new region of the cortex, the angle of actin flow – and hence cell motion – was hardly deflected (Fig. 3K). However, increasing the strength of the perturbation had an increased capacity to reorient the flow (Fig. 3L), suggesting that an external cue could steer cell motion by increasing the strength of actin polymerisation. Indeed, this mechanism of cell steering may be occurring in hemocytes migrating towards wounds as their increase in resultant edge velocity in the direction of motion is likely driven by increased actin polymerisation (Fig. 3G,H). Nevertheless, this minimal model suggests that actin flow organisation is inherently stable and strongly dependent on myosin contraction.

Negatively divergent regions of the actin flowfield represent regions of actin network strain and disassembly

Due to the presence of stable, negatively divergent regions within the actin flowfield, we hypothesised that global actin flow may be coordinated by these points within the network. There are two, non-mutually exclusive mechanisms hypothesised to contribute to actin flows: motor-driven contraction and actin network destruction. We therefore examined how the negatively divergent regions of the network correlated with measures of compression and disassembly. We first calculated the principal component of the strain rate, which is quantified from the spatial changes in the actin velocity field; this analysis of network deformation highlighted that the

negatively divergent regions were correlated with high rates of compression (Fig. 4A,B,D; Supplementary Video 13). We also modelled the assembly/disassembly within the network by taking into account the actin intensity and flow information as previously described¹⁰. This revealed that the negatively divergent regions were also correlated with regions of disassembly (Fig. 4A,C,E; Supplementary Video 13). These data suggest that the negatively divergent regions of the network are controlled by a combination of both contraction and disassembly of actin filaments.

Myosin-II driven contraction and cofilin-mediated disassembly are essential for actin flow

We next examined zygotic mutations in *non-muscle myosin-II* and *cofilin*, which have both been hypothesised to regulate actin flow through contraction and severing, respectively^{10, 41-43}. Indeed, homozygous mutation of either *myosin-II* or *cofilin* led to defects in hemocyte dispersal (Fig. 5A,B; Supplementary Video 14). Furthermore, both mutations showed a reduction in cell speed, and similar to what has been observed in cultured cells *in vitro*, a reduction in actin flow velocity^{10, 41, 42, 44} (Fig. 5C-E; Supplementary Video 15).

We subsequently examined how organisation of the actin flow was affected by the absence of either Cofilin or Myosin-II. As mechanical gradients across the cytoplasm are hypothesised to be a property of polarised motility^{31, 45, 46}, we first determined if there was a gradient of negative divergence. *Wild-type* cells showed a gradient of network divergence starting a few microns from the cell edge, which increased until peaking just before reaching the cell body (Fig. 6A,B). In contrast, in both *myosin-II* and *cofilin* mutant cells, the overall divergence values increased and there was no obvious gradient from front to rear (Fig. 6A,B; Supplementary Video

16). Furthermore, the primary streamline sink, which was negatively divergent in *wild-type* cells, showed an increase in divergence values in the mutants (Fig. 6C) suggesting that myosin-II and cofilin are both playing some role in generating sinks within the actin network.

While the divergence profiles appeared similar in *myosin-II* and *cofilin* mutants, they showed other phenotypes suggestive of unique roles in the regulation of actin flow when we compared rates of assembly/disassembly. *Wild-type* cells displayed a gradient of disassembly that peaked at the rear of the network in a region similar in location to the peak in negative divergence (Fig. 6A,B,D,E). In *cofilin* and *myosin-II* mutant cells, overall disassembly was reduced suggesting that they both play a role in network destruction (Fig. 6D,E; Supplementary Video 16), however, the profile of the disassembly rates was not identical. In the absence of myosin-II, net disassembly of the network was relatively flat until reaching the rear of the network. In contrast, in the absence of cofilin there was a similar profile of net disassembly to *wild-type* cells, with a peak at the rear of the network that failed to reach levels observed in controls (Fig. 6D,E). These data suggest that both cofilin-mediated severing and myosin-II contraction are essential to regulate disassembly, however, cofilin is setting a baseline level of actin depolymerisation across the network while myosin-II is controlling its graded destruction.

Streamline analysis also revealed that the *myosin-II* mutants showed a much more disorganised actin flow. Quantifying the strength of the streamline sink revealed that the maximum streamline endpoint in *myosin-II* mutants accumulated far fewer streamlines than either *wild-type* or *cofilin* mutants (Fig. 6F,G; Supplementary Video 16). Furthermore, quantifying local alignment of the flowfield

revealed that *myosin-II* mutants specifically had a more disorganised actin flow profile (Extended Data Fig. 5A-C).

We also examined how changing polymerisation dynamics altered global actin flow organisation. As loss of actin polymerisation factors (e.g. Scar and Arp2/3) results in a severe and near complete loss of lamellae^{47, 48} it was not possible to analyse their role in controlling actin flow. However, *Drosophila* Ena/Vasp, which enhances leading edge dynamics, plays a more subtle role in regulating hemocyte lamellipodia^{49, 50}, allowing us to examine how changing edge activity affects actin flow. *Ena* mutant hemocytes showed a reduction in edge activity, and consequently a reduction in actin flow speed and an increase in divergence values (Extended Data Fig. 5D-G; Supplementary Video 17). However, we observed no obvious change in actin flow organisation (Extended Data Fig. 5H). Therefore, as predicted by the modelling, the emergence of a stable flow profile is likely insensitive to changes in actin polymerisation.

A gradient of Myosin-II indirectly leads to actin network contraction

As *myosin-II* mutants showed a more perturbed organisation in actin flow, myosin-II may be driving long-range coordination of the network. Indeed, GFP-tagged Myosin-II revealed a front to rear gradient of puncta flowing within the hemocyte lamella (Fig. 7A,B; Supplementary Video 18). We therefore hypothesised that myosin-II may directly control the stable regions of network compression. We simultaneously analysed actin and myosin-II flows while also calculating the divergence within the actin flowfield. To our surprise, we observed no correlation of Myosin-II puncta with divergent hotspots (Fig. 7C; Extended Data Fig. 6A). Indeed, dynamic analysis of the divergence revealed that the divergent hotspots often

developed adjacent to Myosin-II puncta and in between dense actin fibres within the network (Fig. 7D). Furthermore, while the negatively divergent regions appeared to be fixed points within the network, Myosin-II puncta flowed through these regions suggesting that these stable sites of network compression and disassembly are helping drive the flowfield (Supplementary Video 19). Consistent with this, we observed that Myosin-II puncta, while moving in the same direction as the overall actin flow, showed a statistically lower speed, and concomitantly a distinct divergence profile (Extended Data Fig. 6B-E; Supplementary Video 20), which is similar to what was observed in migrating fish keratocytes⁴⁴. These data suggest that myosin-II is not directly responsible for generating local contractile stresses within the actin network sinks; instead, the points of actin network divergence are likely an emergent behaviour driven by a stable gradient of actin network tension and disassembly of the network.

Discussion

Here, we have taken advantage of hemocyte dispersal to examine stereotypical behaviours hypothesised to control motility and determine how these processes correlate with cell directionality. Contrary to the previous lamellipodial-centric model of cell motility, the leading edge is poorly correlated with cell motion with a persistence that is less than the overall persistence of the cell. Hemocytes spend significant energy using extensions to explore their environment rather than directly inducing motion. While this mode of motility has been termed ‘inefficient’^{27, 33}, this does not mean that these seemingly extraneous edge fluctuations are non-functional; indeed, hemocytes are necessary to engulf apoptotic debris⁵¹ and evenly

deposit extracellular matrix⁵², and the decoupling of extensions from motion may be necessary for hemocytes to efficiently explore their environment to carry out these critical tasks. Nevertheless, the term 'leading edge' is a misnomer as it is not obviously playing a leading role during hemocyte random migration; however, nor is it completely uncorrelated with motion. Edge fluctuations in hemocytes are still weakly correlated to motion, and therefore cells must have an intrinsic capacity to integrate this activity to 'decide' on a direction of travel, which we speculate occurs through the flow of actin. Furthermore, edge extensions, actin flow, and cell motion are highly integrated behaviours with no obvious temporal hierarchy. The migratory process is not stepwise and, in the future, only a holistic approach to understanding motility may explain how these behaviours are coordinated in such a precise fashion to control coherent cell motion.

Our global view of actin flow revealed a structure that is coordinated across the entire cell, both for hemocytes and other cell types. Indeed, this is consistent with what was reported in one of the first publications of actin flow³⁴; it is also interesting to note that the authors of this work presciently noted that organisation of the flow, in contrast to the leading edge, "is time persistent over minutes", and we hypothesise that this stable organisation of the actin flowfield may be a consistent feature of motility. The network sink also represents a transition from retrograde to anterograde actin flow. While actin flows within migrating cells are often generally termed 'retrograde flow' (due to the focus on the leading edge), there is significant anterograde motion observed from the rear of numerous cell types^{18, 34-36}. Furthermore, modelling has predicted that retrograde flow at the front of a migrating cell will transition to anterograde flow as the adhesions switch from gripping (within the retrograde region) to slipping (within the anterograde region)^{53, 54}.

One outstanding question is, what is controlling the formation and stability of these network sinks? As the motion of the leading edge and the sinks are correlated, it is possible that there is information being transmitted between these two sites. Recent work has suggested that actin flow mediates a coupling between cell speed and persistence through the advection of polarity cues from the leading edge³¹ and it is possible that these cues may converge on the network sink. Another possibility is that the flowing actin network is inherently stable and flow patterns may develop spontaneously in the absence of any direct regulation. Indeed, our minimal model of actin flow, in which flows emerge primarily through myosin-II contraction, leads to a highly stable 'sink' at the rear.

The destruction of the actin network, which is occurring at these network sinks may also be directly providing forces for locomotion. Disassembly of cytoskeletal networks can generate force in the absence of motors through entropic contraction⁵⁵⁻⁵⁷. Due to the absence of a time lag between the direction of nuclear movement and the primary streamline sink, we hypothesise that the sink may provide the force for motion of the trailing cell body. The actin network on the retrograde side of this sink is experiencing high friction while the anterograde side is slipping. This would imply that the retrograde region is anchored to the substrate, allowing the forces generated by the reorganisation of the actin network at the sink to drive unidirectional retraction of the rear of the cell. This mechanism is also consistent with the network-contraction model that has been hypothesised to drive rear retraction in other cell types⁵⁸.

This organisation of actin flow may have wide-ranging implications for how cells interpret and respond to cues. Due to the extreme stability of actin flow, it is possible that in some cells a complete loss of polarity may be required to reset flow direction and redirect cell motion (e.g. during run and tumble modes of migration⁵⁹).

406 The stable actin flow may also be providing a stable polarity to the cell that enhances
407 the discrimination of guidance cues. While internal amplification through reaction-
408 diffusion signalling modules are hypothesised to be required to accurately chemotax
409 towards low concentrations of guidance cues^{4, 60}, this may be unnecessary. The
410 stable flow of actin itself may be sufficient to provide the directional memory that
411 allows the leading edge to rapidly sample external cues with subtle biases in the
412 edge fluctuations stabilised and integrated by the actin flow. Indeed, directional
413 memory can make chemotaxis more efficient and discriminatory^{61, 62}, and while
414 reaction-diffusion modules have been hypothesised to control this memory⁶³, recent
415 work suggests that it may also come from the cytoskeleton itself⁶⁴; our work
416 suggests that it may originate from the highly coordinated and stable flow of the actin
417 network.

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Figure Legends

Figure 1. Leading edge fluctuations are a weak predictor of cell directionality

(A-E) Morphodynamic analysis of edge fluctuation in hemocytes by quantifying the speed of: (A) all edge extensions, (B) the maximum edge extension (longest contiguous extension of perimeter), (C) extensions in the direction of motion (30° cone), (D) edge extensions and retractions in the direction of motion, and (E) all edge extensions and retractions (green, extension; magenta, retraction; for display purposes 'A-E' show unit vectors; scale bar 10 μm).

(F) Comparison of cell speed with the speed of edge fluctuations as measured in 'A-E'. *** $P < 0.0001$, Kruskal-Wallis and Dunn's multiple comparison test. Boxplot shows medians, 25th and 75th percentiles as box limits, 10th and 90th as whiskers ($n = 443$, 9 biologically independent samples). Note that cell speed is not correlated with edge extension speed, and that net extensions (extensions – retractions) sum to zero showing that cells maintain a constant area over time.

(G) A representative snapshot of a randomly migrating hemocyte with edge extensions automatically segmented. Vectors are drawn from the nucleus to each individual extension (blue arrows) and correlated with the direction of cell motion (magenta arrow) in panel 'H'. Scale bar 10 μm .

(H) A rose plot showing the direction of all extension vectors, as highlighted in 'G', normalised to the direction of cell motion ($n = 16379$, 9 biologically independent samples).

(I) A representative snapshot of a randomly migrating hemocyte with the maximum edge extension (by area) automatically segmented. Vectors are drawn from the nucleus to the centroid of the maximum extension (green arrow) and correlated with the direction of cell motion (magenta arrow) in 'J'.

(J) A rose plot showing the direction of the maximum extension vectors, as highlighted in 'I', normalised to the direction of cell motion ($n = 443$, 9 biologically independent samples).

(K) Directional autocorrelation comparing the persistence of cell motion and maximum edge extension showing that the maximum edge extension is less persistent than overall cell motion (note that a slower decay represents an increased persistence). Dotted lines are real data and solid lines represent fitted decay curves ($n = 9$ biologically independent samples).

Figure 2. Actin retrograde flow is globally organised in migrating hemocytes

(A) Particle Image Velocimetry (PIV) analysis performed on a LifeAct-GFP expressing cell to highlight the direction and magnitude of actin flow. The region of the flowfield without vectors represents the soma of the hemocyte, which has no observable actin flow, and this information was removed for all subsequent quantification. Scale bar 10 μm .

(B) Divergence calculated from the actin flowfield to highlight sinks within the network. In this image only negatively divergent regions are highlighted.

(C) Streamlines calculated from the actin flowfield in which streamlines were seeded along the boundary of the cell.

(D) The confluence of streamlines quantified by calculating the number of streamlines ending in any location within the cell. In this image, the size of the spot is normalised to the number of streamline endpoints.

(E) The actin flow divergence calculated at the primary sink and compared to the divergence values averaged across the entire cell. *** $P < 0.0001$, Mann-Whitney two-tailed test. Boxplot shows medians, 25th and 75th percentiles as box limits, 10th and 90th as whiskers ($n = 443$, 9 biologically independent samples).

(F) A correlation map in which the direction of cell motion was correlated to the direction of every actin flow vector within the cell. Note that a positive correlation highlights anterograde flow while a negative correlation denotes retrograde flow. The dashed circle indicates the location of the primary sink in this frame of the movie.

(G) Quantification of the gradient of the correlation map in 'F' reveals sharp transition regions within the flowfield. The dashed circle indicates the location of the primary streamline sink in this frame of the time-lapse movie.

(H) Quantification of the gradient of the retrograde/anterograde flow correlation, as highlighted in 'F' and 'G', at the primary streamline sinks compared to the gradient values averaged across the entire cell. *** $P < 0.0001$, Mann-Whitney two-tailed test. Boxplot representation as in 'E' ($n = 443$, 9 biologically independent samples).

Figure 3. The polarity of global actin flow is highly stable and correlated to hemocyte motion during random and directed migration

(A) Schematic depicting the regions correlated to cell motion. *** $P < 0.0001$, (n.s) $P = 0.4471$, Kruskal-Wallis and Dunn's multiple comparison test. Boxplot shows medians, 25th and 75th percentiles as box limits, 10th and 90th as whiskers ($n = 443$, 9 biologically independent samples).

(B) Directional autocorrelation of behaviours during random migration. Dotted lines, real data; solid lines, fitted decay curves ($n = 9$ biologically independent samples, same data as in '1K').

(C) Temporal cross-correlation of migratory behaviours revealing a peak at 0-lag ($n = 9$ biologically independent samples).

(D) Visualisation of all hemocyte edge extensions (green) during random and directed migration.

(E) Schematic of a directly migrating cell correlating behaviours as in 'A'.

(F) Directional autocorrelation of behaviours during directed migration. Dotted lines, real data; solid lines, fitted decay curves ($n = 4$ biologically independent samples).

(G) Correlation to motion of edge extension unit vectors (i.e. direction only) during random ($n = 33617$, 9 biologically independent samples) versus directed migration ($n = 19486$, 4 biologically independent samples). (n.s) $P = 0.4155$, Mann-Whitney two-tailed test. Boxplot representation as in 'A'.

(H) Correlation to motion of the resultant edge extension velocities (i.e. direction and magnitude) during random ($n = 443$, 9 biologically independent samples) versus

directed migration ($n = 272$, 4 biologically independent samples). *** $P = 0.0002$, Mann-Whitney two-tailed test. Boxplot representation as in 'A'.

(I) A minimal one-dimensional fluid-mechanical model shows myosin contraction spontaneously leads to acto-myosin cortical flows with a sink at the rear ($\theta = 0$), corresponding with a peak in Myosin-II concentration.

(J) Linear stability analysis shows the emergence of flows is sensitive to Myosin-II levels (steep dependence in path 2) and insensitive to polymerisation rate (flat dependence in path 1).

(K) Angular location of Myosin-II peak and hence direction of cell migration (red solid line) versus the angle of actin perturbation (black dashed line).

(L) Angular location of Myosin-II peak and cell motion versus the strength of actin perturbation. Angle of perturbation (black dashed line) was kept constant while varying its strength.

Figure 4. Negatively divergent regions of the actin flowfield represent regions of actin network strain and disassembly

(A-C) Heatmaps comparing the quantification of divergence (A), network compression (B), and actin network disassembly (C) in an individual hemocyte. Bottom panels are high magnification images of the boxes outlined in the upper panels. Note the partial overlap of these parameters.

(D) Scatter plot comparing a random sample of points in the actin flowfield for divergence and principal strain. Note the positive relationship between negative divergence and the negative values of the principal strain (i.e. compression) ($n = 5000$ random points, 9 biologically independent samples).

(E) Scatter plot comparing a random sample of points in the actin flowfield for divergence and amount of assembly/disassembly within the actin network. Note the positive relationship between negative divergence and the amount of disassembly ($n = 5000$ random points, 9 biologically independent samples).

Figure 5. Loss of myosin-II and cofilin lead to reduced actin flow and cell speed

(A) Images of hemocytes on the ventral surface of *Drosophila* embryos in *wild-type*, *myosin-II*, and *cofilin* mutant embryos. LifeAct-GFP is shown in green, nuclei are labelled in magenta. Scale bar 30 μm .

(B) Temporal average projection of *wild-type*, *myosin-II*, and *cofilin* mutant embryos, highlighting domains occupied by migrating hemocytes. Note that the mutant embryos display a less homogenous domain distribution. Scale bar 30 μ m.

(C) PIV analysis of actin flow in *wild-type*, *myosin-II*, and *cofilin* mutant cells.

(D) Quantification of mean cell speed in *wild-type* and mutant cells showing that both *myosin-II* and *cofilin* mutant hemocytes are slower than *wild-type* cells. ***P < 0.001, ordinary one-way ANOVA test and Holm-Sidak's multiple comparison test. Boxplot shows medians, 25th and 75th percentiles as box limits, minimum and maximum values as whiskers; each datapoint is displayed as a dot (n = 9 biologically independent samples for each genotype).

(E) Quantification of mean actin flow speed in *wild-type* and mutant cells. Both *myosin-II* and *cofilin* mutant hemocytes are slower than *wild-type* cells. ***P < 0.001, ordinary one-way ANOVA test and Holm-Sidak's multiple comparison test. Boxplot representation as in 'D' (n = 9 biologically independent samples for each genotype).

Figure 6. A gradient of myosin-II driven contraction is essential for global organisation of actin flow

(A) Heatmaps comparing the quantification of divergence in *wild-type*, *myosin-II*, and *cofilin* mutant hemocytes.

(B) Quantification of the mean divergence values in *wild-type* and mutant cells calculated by drawing linescans from the cell body to the edge (*wild-type*, n = 1329 lines, 9 biologically independent samples; *myosin-II* mutant, n = 657 lines, 4 biologically independent samples; *cofilin* mutant, n = 1480 lines, 7 biologically independent samples). Error bars = SEM.

(C) Quantification of the mean divergence at primary sink in *wild-type* and mutant cells, which reveals that mutants have an increase in divergence values highlighting a reduction in network compression. ***P = 0.0002, **P = 0.004, Kruskal-Wallis and Dunn's multiple comparison test. Boxplot shows medians, 25th and 75th percentiles as box limits, minimum and maximum values as whiskers; each datapoint is displayed as a dot (n = 9 biologically independent samples for all genotypes).

(D) Heatmaps comparing the quantification of actin disassembly in *wild-type* and mutant cell

(E) Quantification of the mean assembly/disassembly values in *wild-type* and mutant cells calculated by drawing linescans from the cell body to the edge (*wild-type*, n =

1328 lines, 9 biologically independent samples; *myosin-II* mutant, n = 657 lines, 4 biologically independent samples; *cofilin* mutant, n = 1538 lines, 7 biologically independent samples). Error bars = SEM.

(F) Images highlighting an analysis of streamlines through the actin flowfield in *wild-type* and mutant cells. Note the disorganised streamlines in *myosin-II* mutants.

(G) Quantification of the percentage of streamlines that end at the primary sink in *wild-type* and mutant cells. Note that *myosin-II* mutants show a significant reduction in their streamline confluence compared to *wild-type* or *cofilin* mutant cells. **P = 0.0018, (n.s) P = 0.6840, Kruskal-Wallis test and Dunn's multiple comparison test. Boxplot representation as in 'C' (n = 9 biologically independent samples for each genotype).

Figure 7. A gradient of myosin-II indirectly leads to actin network contraction

(A) Temporal maximum projection of a Myosin-II-GFP expressing hemocyte highlighting that Myosin-II puncta within the lamellae are predominantly toward the rear of the network surrounding the cell body. Scale bar 10 μ m.

(B) Linescan analysis of Actin and Myosin-II localisation within hemocytes. The plot profile of mean fluorescence intensity was performed on randomly chosen lines within the lamellae from the cell body towards the edge (see insert). Note that the average intensity of Myosin-II is high toward the cell body and decreases in a gradient approaching the edge, whereas Actin intensity remains constant (n = 63 lines, 12 independent samples). Error bars = SEM.

(C) Comparison of Actin and Myosin-II localisation with actin flow divergence. Bottom panels are high magnification images of the boxes outlined in upper panels. Asterisks highlight regions of strong negative divergence, which show no obvious colocalisation with Myosin-II. Scale bar 10 μ m.

(D) Time-lapse series comparing Actin and Myosin-II localisation with actin flow divergence. Circles highlight example regions of strong negative divergence. Note that the negatively divergent regions are adjacent to Myosin-II puncta.

615 References

- 616 1. Ridley, A.J. *et al.* Cell migration: integrating signals from front to back.
617 *Science* **302**, 1704-1709 (2003).
- 618 2. Diz-Munoz, A., Fletcher, D.A. & Weiner, O.D. Use the force: membrane
619 tension as an organizer of cell shape and motility. *Trends Cell Biol* **23**, 47-53
620 (2013).
- 621 3. Shi, C., Huang, C.H., Devreotes, P.N. & Iglesias, P.A. Interaction of motility,
622 directional sensing, and polarity modules recreates the behaviors of
623 chemotaxing cells. *PLoS Comput Biol* **9**, e1003122 (2013).
- 624 4. Xiong, Y., Huang, C.H., Iglesias, P.A. & Devreotes, P.N. Cells navigate with a
625 local-excitation, global-inhibition-biased excitable network. *Proc Natl Acad Sci*
626 *U S A* **107**, 17079-17086 (2010).
- 627 5. Houk, A.R. *et al.* Membrane tension maintains cell polarity by confining
628 signals to the leading edge during neutrophil migration. *Cell* **148**, 175-188
629 (2012).
- 630 6. Gorelik, R. & Gautreau, A. Quantitative and unbiased analysis of directional
631 persistence in cell migration. *Nat Protoc* **9**, 1931-1943 (2014).
- 632 7. Machesky, L.M. & Insall, R.H. Scar1 and the related Wiskott-Aldrich syndrome
633 protein, WASP, regulate the actin cytoskeleton through the Arp2/3 complex.
634 *Curr Biol* **8**, 1347-1356 (1998).
- 635 8. Welch, M.D., DePace, A.H., Verma, S., Iwamatsu, A. & Mitchison, T.J. The
636 human Arp2/3 complex is composed of evolutionarily conserved subunits and
637 is localized to cellular regions of dynamic actin filament assembly. *J Cell Biol*
638 **138**, 375-384 (1997).
- 639 9. Lin, C.H., Espreafico, E.M., Mooseker, M.S. & Forscher, P. Myosin drives
640 retrograde F-actin flow in neuronal growth cones. *Biol Bull* **192**, 183-185
641 (1997).
- 642 10. Wilson, C.A. *et al.* Myosin II contributes to cell-scale actin network treadmilling
643 through network disassembly. *Nature* **465**, 373-377 (2010).
- 644 11. Ponti, A., Machacek, M., Gupton, S.L., Waterman-Storer, C.M. & Danuser, G.
645 Two distinct actin networks drive the protrusion of migrating cells. *Science*
646 **305**, 1782-1786 (2004).
- 647 12. Hu, K., Ji, L., Applegate, K.T., Danuser, G. & Waterman-Storer, C.M.
648 Differential transmission of actin motion within focal adhesions. *Science* **315**,
649 111-115 (2007).
- 650 13. Gardel, M.L. *et al.* Traction stress in focal adhesions correlates biphasically
651 with actin retrograde flow speed. *J Cell Biol* **183**, 999-1005 (2008).
- 652 14. Ballestrem, C., Hinz, B., Imhof, B.A. & Wehrle-Haller, B. Marching at the front
653 and dragging behind: differential α V β 3-integrin turnover regulates focal
654 adhesion behavior. *J Cell Biol* **155**, 1319-1332 (2001).
- 655 15. Mitchison, T.J. & Cramer, L.P. Actin-based cell motility and cell locomotion.
656 *Cell* **84**, 371-379 (1996).
- 657 16. Krause, M. & Gautreau, A. Steering cell migration: lamellipodium dynamics
658 and the regulation of directional persistence. *Nat Rev Mol Cell Biol* **15**, 577-
659 590 (2014).
- 660 17. Wu, C. *et al.* Arp2/3 is critical for lamellipodia and response to extracellular
661 matrix cues but is dispensable for chemotaxis. *Cell* **148**, 973-987 (2012).

- 662 18. Gupton, S.L. *et al.* Cell migration without a lamellipodium: translation of actin
663 dynamics into cell movement mediated by tropomyosin. *J Cell Biol* **168**, 619-
664 631 (2005).
- 665 19. Fritz-Laylin, L.K. *et al.* Actin-based protrusions of migrating neutrophils are
666 intrinsically lamellar and facilitate direction changes. *Elife* **6** (2017).
- 667 20. Leithner, A. *et al.* Diversified actin protrusions promote environmental
668 exploration but are dispensable for locomotion of leukocytes. *Nat Cell Biol* **18**,
669 1253-1259 (2016).
- 670 21. Keren, K. *et al.* Mechanism of shape determination in motile cells. *Nature* **453**,
671 475-480 (2008).
- 672 22. Tweedy, L., Meier, B., Stephan, J., Heinrich, D. & Endres, R.G. Distinct cell
673 shapes determine accurate chemotaxis. *Sci Rep* **3**, 2606 (2013).
- 674 23. Vallotton, P. & Small, J.V. Shifting views on the leading role of the
675 lamellipodium in cell migration: speckle tracking revisited. *J Cell Sci* **122**,
676 1955-1958 (2009).
- 677 24. Abercrombie, M. Croonian Lecture, 1978 - Crawling Movement of Metazoan
678 Cells. *Proc R Soc Ser B-Bio* **207**, 129-+ (1980).
- 679 25. Danuser, G., Allard, J. & Mogilner, A. Mathematical modeling of eukaryotic
680 cell migration: insights beyond experiments. *Annu Rev Cell Dev Biol* **29**, 501-
681 528 (2013).
- 682 26. Machacek, M. & Danuser, G. Morphodynamic profiling of protrusion
683 phenotypes. *Biophys J* **90**, 1439-1452 (2006).
- 684 27. Hermans, T.M. *et al.* Motility efficiency and spatiotemporal synchronization in
685 non-metastatic vs. metastatic breast cancer cells. *Integr Biol (Camb)* **5**, 1464-
686 1473 (2013).
- 687 28. Betz, T., Koch, D., Lim, D. & Kas, J.A. Stochastic actin polymerization and
688 steady retrograde flow determine growth cone advancement. *Biophys J* **96**,
689 5130-5138 (2009).
- 690 29. Davis, J. *et al.* Emergence of embryonic pattern through contact inhibition of
691 locomotion. *Development* **139**, 4555-4560 (2012).
- 692 30. Davis, J.R. *et al.* Inter-cellular forces orchestrate contact inhibition of
693 locomotion. *Cell* **161**, 361-373 (2015).
- 694 31. Maiuri, P. *et al.* Actin flows mediate a universal coupling between cell speed
695 and cell persistence. *Cell* **161**, 374-386 (2015).
- 696 32. Wood, W., Faria, C. & Jacinto, A. Distinct mechanisms regulate hemocyte
697 chemotaxis during development and wound healing in *Drosophila*
698 *melanogaster*. *J Cell Biol* **173**, 405-416 (2006).
- 699 33. Dunn, G., Weber, I. & Zicha, D. Protrusion, retraction and the efficiency of cell
700 locomotion. *Math Biosci Interac*, 33-46 (1997).
- 701 34. Vallotton, P., Gupton, S.L., Waterman-Storer, C.M. & Danuser, G.
702 Simultaneous mapping of filamentous actin flow and turnover in migrating
703 cells by quantitative fluorescent speckle microscopy. *Proc Natl Acad Sci U S*
704 *A* **101**, 9660-9665 (2004).
- 705 35. Gupton, S.L. & Waterman-Storer, C.M. Spatiotemporal feedback between
706 actomyosin and focal-adhesion systems optimizes rapid cell migration. *Cell*
707 **125**, 1361-1374 (2006).
- 708 36. Fournier, M.F., Sauser, R., Ambrosi, D., Meister, J.J. & Verkhovsky, A.B.
709 Force transmission in migrating cells. *J Cell Biol* **188**, 287-297 (2010).
- 710 37. Betz, T., Koch, D., Lu, Y.B., Franze, K. & Kas, J.A. Growth cones as soft and
711 weak force generators. *Proc Natl Acad Sci U S A* **108**, 13420-13425 (2011).

- 712 38. Razzell, W., Evans, I.R., Martin, P. & Wood, W. Calcium flashes orchestrate
713 the wound inflammatory response through DUOX activation and hydrogen
714 peroxide release. *Curr Biol* **23**, 424-429 (2013).
- 715 39. Hawkins, R.J. *et al.* Spontaneous contractility-mediated cortical flow
716 generates cell migration in three-dimensional environments. *Biophys J* **101**,
717 1041-1045 (2011).
- 718 40. Bois, J.S., Julicher, F. & Grill, S.W. Pattern formation in active fluids. *Phys*
719 *Rev Lett* **106**, 028103 (2011).
- 720 41. Vitriol, E.A., Wise, A.L., Berginski, M.E., Bamburg, J.R. & Zheng, J.Q.
721 Instantaneous inactivation of cofilin reveals its function of F-actin disassembly
722 in lamellipodia. *Mol Biol Cell* **24**, 2238-2247 (2013).
- 723 42. Delorme, V. *et al.* Cofilin activity downstream of Pak1 regulates cell protrusion
724 efficiency by organizing lamellipodium and lamella actin networks. *Dev Cell*
725 **13**, 646-662 (2007).
- 726 43. Medeiros, N.A., Burnette, D.T. & Forscher, P. Myosin II functions in actin-
727 bundle turnover in neuronal growth cones. *Nat Cell Biol* **8**, 215-226 (2006).
- 728 44. Schaub, S., Bohnet, S., Laurent, V.M., Meister, J.J. & Verkhovsky, A.B.
729 Comparative maps of motion and assembly of filamentous actin and myosin II
730 in migrating cells. *Mol Biol Cell* **18**, 3723-3732 (2007).
- 731 45. Lomakin, A.J. *et al.* Competition for actin between two distinct F-actin
732 networks defines a bistable switch for cell polarization. *Nat Cell Biol* **17**, 1435-
733 1445 (2015).
- 734 46. Taylor, D.L. The contractile basis of amoeboid movement. IV. The
735 viscoelasticity and contractility of amoeba cytoplasm in vivo. *Exp Cell Res*
736 **105**, 413-426 (1977).
- 737 47. Evans, I.R., Ghai, P.A., Urbancic, V., Tan, K.L. & Wood, W. SCAR/WAVE-
738 mediated processing of engulfed apoptotic corpses is essential for effective
739 macrophage migration in *Drosophila*. *Cell Death Differ* **20**, 709-720 (2013).
- 740 48. Verboon, J.M., Rahe, T.K., Rodriguez-Mesa, E. & Parkhurst, S.M. Wash
741 functions downstream of Rho1 GTPase in a subset of *Drosophila* immune cell
742 developmental migrations. *Mol Biol Cell* **26**, 1665-1674 (2015).
- 743 49. Tucker, P.K., Evans, I.R. & Wood, W. Ena drives invasive macrophage
744 migration in *Drosophila* embryos. *Dis Model Mech* **4**, 126-134 (2011).
- 745 50. Davidson, A.J., Millard, T.H., Evans, I.R. & Wood, W. Ena orchestrates
746 remodelling within the actin cytoskeleton to drive robust *Drosophila*
747 macrophage chemotaxis. *J Cell Sci* **132** (2019).
- 748 51. Zhou, L., Hashimi, H., Schwartz, L.M. & Nambu, J.R. Programmed cell death
749 in the *Drosophila* central nervous system midline. *Curr Biol* **5**, 784-790 (1995).
- 750 52. Matsubayashi, Y. *et al.* A Moving Source of Matrix Components Is Essential
751 for De Novo Basement Membrane Formation. *Curr Biol* **27**, 3526-3534 e3524
752 (2017).
- 753 53. Shao, D., Levine, H. & Rappel, W.J. Coupling actin flow, adhesion, and
754 morphology in a computational cell motility model. *Proc Natl Acad Sci U S A*
755 **109**, 6851-6856 (2012).
- 756 54. Fuhs, T., Goegler, M., Brunner, C.A., Wolgemuth, C.W. & Kaes, J.A. Causes
757 of retrograde flow in fish keratocytes. *Cytoskeleton (Hoboken)* **71**, 24-35
758 (2014).
- 759 55. Miao, L., Vanderlinde, O., Stewart, M. & Roberts, T.M. Retraction in amoeboid
760 cell motility powered by cytoskeletal dynamics. *Science* **302**, 1405-1407
761 (2003).

56. Zajac, M., Dacanay, B., Mohler, W.A. & Wolgemuth, C.W. Depolymerization-driven flow in nematode spermatozoa relates crawling speed to size and shape. *Biophys J* **94**, 3810-3823 (2008).
57. Mseka, T. & Cramer, L.P. Actin depolymerization-based force retracts the cell rear in polarizing and migrating cells. *Curr Biol* **21**, 2085-2091 (2011).
58. Svitkina, T.M., Verkhovsky, A.B., McQuade, K.M. & Borisy, G.G. Analysis of the actin-myosin II system in fish epidermal keratocytes: mechanism of cell body translocation. *J Cell Biol* **139**, 397-415 (1997).
59. Reichman-Fried, M., Minina, S. & Raz, E. Autonomous modes of behavior in primordial germ cell migration. *Dev Cell* **6**, 589-596 (2004).
60. Iglesias, P.A. & Devreotes, P.N. Biased excitable networks: how cells direct motion in response to gradients. *Curr Opin Cell Biol* **24**, 245-253 (2012).
61. Skoge, M. *et al.* Cellular memory in eukaryotic chemotaxis. *Proc Natl Acad Sci U S A* **111**, 14448-14453 (2014).
62. Aquino, G., Tweedy, L., Heinrich, D. & Endres, R.G. Memory improves precision of cell sensing in fluctuating environments. *Sci Rep* **4**, 5688 (2014).
63. Nakajima, A., Ishihara, S., Imoto, D. & Sawai, S. Rectified directional sensing in long-range cell migration. *Nat Commun* **5**, 5367 (2014).
64. Prentice-Mott, H.V. *et al.* Directional memory arises from long-lived cytoskeletal asymmetries in polarized chemotactic cells. *Proc Natl Acad Sci U S A* **113**, 1267-1272 (2016).

Materials and methods

Fly genetics

The following fly stocks were used in this study: *w¹¹¹⁸* strain as *wild-type* (Bloomington Drosophila Stock Center (BDSC), BL3605); *myosin-II* mutant (BDSC, BL4199); *cofilin* mutant (BDSC, BL9107), *ena* mutant (BDSC, BL8569). Hemocytes were labelled using the promoters, *Srp-Gal4* (BDSC7, BL78565) or *Sn-Gal4*⁶⁵. The following fluorescent probes were used to label: nuclei [UAS-RedStinger, (BDSC, BL8546 and 8547)]; actin [UAS-LifeAct-GFP⁶⁵ or UAS-Moesin-Cherry⁶⁶, or UAS-Act5C-GFP (BDSC, BL9275)]; Myosin-II heavy chain [(UAS-Zip-GFP)⁶⁷]. Flies were left to lay eggs on grape juice/agar plates overnight at 25°C. Embryos were dechorionated in bleach and the appropriate genotype was identified based on the presence of fluorescent markers.

Cell lines

Zebrafish lines expressing LifeAct-GFP (Tg(actb1:lifeact-GFP)) were generated and fish keratocytes cultured as previously described⁶⁸. Keratocytes were prepared from adult zebrafish scales, plucked from sacrificed animals and washed three times with Dulbecco's Modified Eagle's Medium (DMEM) (Gibco). Scales were incubated in START medium at room temperature for one or two days to allow the keratocytes to migrate off in a monolayer. The monolayer was then washed three times in PBS, incubated for 40 min with Running Buffer with 1 mM EGTA and the scales removed. The remaining cells were washed three times with PBS, trypsinised for 2 min with 0.25% Trypsin-EDTA (Gibco) at room temperature, resuspended in the same volume of trypsin inhibitor (Sigma) and transferred to a coverslip coated with 0.5mg/ml PLL(20)-g[3.5]-PEG(2)/PEG(3.4)-RGD (Surface Solutions) for 50min. Live cell imaging of migrating keratocytes was performed at room temperature in START medium. Confocal microscopy was performed with an inverted microscope (Zeiss), equipped with a Spinning disk system (Yokogawa X1, iXon897, Andor), a C-Apochromat 100x/1.4 Oil Objective, a motorised stage and a 488nm laser.

RPE1 cells expressing LifeAct-TagRFP (a gift from Buzz Baum)⁶⁹ were cultured in DMEM/F12 media containing HEPES and sodium bicarbonate (Sigma)

supplemented with 10% Fetal Bovine Serum (HyClone, Fisher Scientific), 1% penicillin-streptomycin (Sigma) and 2 mM L-glutamine (Sigma). Imaging was carried out in 35 mm dishes that had been coated with 10µg/ml fibronectin for 1 hour at 37 °C. RPE1 cells were plated and allowed to adhere and spread overnight before imaging. Cells were imaged using an LSM 880 confocal microscope using airyscan with a 40x NA 1.3 Plan-Apochromat oil objective at 1.8x zoom. Images were acquired every 30 seconds. Nuclei were labelled with SiR-DNA (Spirochrome) at 0.5 mM to enable cell tracking.

Embryo microscopy

Embryos were mounted as previously described⁴ and time-lapse images were acquired every 5 s with a PerkinElmer Ultraview spinning disk microscope equipped with a 63x NA 1.4 Plan-Apochromat oil objective during developmental dispersal (stages 15–16). Whole embryo snapshots were taken using the LSM 880 confocal microscope (Carl Zeiss) equipped with a 40x NA 1.3 Plan-Apochromat oil objective.

Data analysis

For the characterisation of control hemocyte migratory parameters (i.e. directional autocorrelation, retrograde flow speed, streamline analysis, divergence, principal strain, assembly/disassembly, and flow alignment) data were gathered on a per/frame basis from 9 individual cells each imaged over approximately 4-5 minutes at intervals of 5 s/frame, representing an n number of 443 time-points. When comparing control and mutant genotypes, statistical tests were performed on a per cell basis due to the partial penetrance of the various mutant phenotypes. Here, comparisons were made between the 9 control cells, 9 *myosin-II* mutants, and 9 *cofilin* mutants. The specific statistical test, as well as the thresholds for significance are noted in the respective figure legends. The raw data that support the findings of this study are available from the corresponding author upon reasonable request [BS]. The computational analysis was performed in MATLAB (Mathworks®) using custom code, which can be obtained from the corresponding author upon reasonable request [BS].

Wounding

67 Laser wounding was performed using an ablation laser (MicroPoint; Andor
68 Technology) as previously described⁷⁰ and imaged using a Perkin Elmer spinning
69 disk microscope.

70

71 **Photobleaching**

72 Photobleaching experiments were performed on hemocytes labelled with Actin-GFP.
73 Images were acquired every 5 s using airyscan imaging on an LSM 880 confocal
74 microscope (Carl Zeiss), equipped with a 63x NA 1.4 Plan-Apochromat oil objective
75 and a 1.8X zoom. Laser power was set at 100% to bleach a region of 25 x 25 pixels
76 for 2 s, with a pixel dwell time of 66 μ s.

77

78 **Quantification of Moesin-Cherry and Myosin-GFP fluorescence**

79 In order to quantify the spatial distribution of Actin and Myosin-II within hemocytes,
80 cells expressing Moesin-Cherry and Myosin-II-GFP were sampled by linescan
81 analysis to measure average fluorescence intensity. Fiji *line* and *profile* functions
82 were used to draw and record 2 μ m wide lines from the cell body boundary to the
83 cell edge in 12 different cells.

84

85 **Cell tracking**

86 Hemocytes containing labelled nuclei were first thresholded in Fiji. Tracking was then
87 performed in MATLAB by calculating the positions of the centroid of the nucleus
88 through time.

89

90 **Particle Image Velocimetry**

91 Time-lapse images of freely moving hemocytes were acquired at 5 s/frame. Actin
92 was labelled with LifeAct-GFP for all figures with the exception of Fig. 7 and
93 Extended Data Fig. 6, which used Moesin-Cherry in conjunction with Myosin-II-GFP
94 (*Drosophila* non-muscle Myosin heavy chain) Cells were then manually segmented
95 prior to PIV analysis.

96

97 There is no grossly observable actin flow behaviour within the cell body of the
98 hemocytes, therefore information from the cell body was removed by manually
99 segmenting the cell body region and using this as a mask to remove PIV values. The
100 actin flow PIV information is therefore entirely from within the lamellae. To observe

myosin-II flow in the lamellae, the signal from the cell body was oversaturated. For this reason, no PIV information could be obtained from the cell body region and it was excluded from the myosin flow PIV analysis and its actin flow counterpart (Extended Data Fig. 6D-G).

A 2D cross-correlation algorithm adapted from classical PIV was implemented³⁰. In brief, this method compares a region of interest in an image (source image) with a larger region of a subsequent image (search image). The sizes of the source and search regions are determined on the basis of the feature size to be tracked and the area of their expected displacement (i.e. actin bundles). For this analysis, source and search images encompassing areas of $1.2 \mu\text{m}^2$ and $2 \mu\text{m}^2$ were used. A cross-correlation map was computed by analysing the cross-correlation coefficient between the source image and the search image, by shifting the source across the search one pixel at a time. Network displacement was measured by finding the maximum coefficient within the resulting cross-correlation map. To filter anomalous tracking data, only displacements that had a cross-correlation coefficient above a certain threshold, c_0 , were kept. For the present work, the threshold was set at $c_0 = 0.5$. Finally, a spatial convolution with a Gaussian kernel (size of $5 \mu\text{m}$, sigma of $1 \mu\text{m}$), and temporal convolution with temporal kernel of 20 s (sigma 10 s) were used to interpolate the measured displacements to cover all the pixels within the cell outline. The complete algorithm for this analysis was implemented in MATLAB.

Defining retrograde and anterograde flow regions

Retrograde and anterograde flow were defined with respect to their respective alignment to cell motion. The direction of the actin flow at every point within the lamellae was correlated with the instantaneous direction of cell motion using the cosine of the angle between these velocity vectors. Retrograde flow was defined as a negative correlation while anterograde flow was a positive correlation to cell motion.

Streamlines

Streamlines were used to assess the global organisation of the actin flowfield. Here each line is drawn tangent to a local velocity vector and describes a path that a massless particle would take if entering the actin flowfield at that point (MATLAB

stream2 function). The seed points for the streamlines were placed at every pixel along the cell boundary. For visualisation purposes streamlines were represented at regular intervals (MATLAB *streamslice* function). Line Integral Convolution was employed to represent global streamline activity (Figure S2D) utilising an open source vector field visualisation toolkit (<http://sccn.ucsd.edu/~nima/>). Streamline sinks were defined by quantifying the frequency of streamline termini within non-overlapping $2.5 \mu\text{m}^2$ regions of the cell image and the coordinates of these endpoints were set at the centre of the boxes. For position vector analysis of these endpoints, vectors were constructed from the centroid of the nucleus to the endpoint coordinates.

Principal strain

Local deformation of the actin network can be quantified by evaluation of the principal strains which are derived from local velocity changes obtained by PIV. The relative positional changes of points within a deforming body are described with a velocity tensor, which is computed based on a central difference estimation over $2.5 \mu\text{m}$ in both spatial dimensions.

$$V = \begin{pmatrix} \frac{\partial V_x}{\partial x} & \frac{\partial V_x}{\partial y} \\ \frac{\partial V_y}{\partial x} & \frac{\partial V_y}{\partial y} \end{pmatrix}$$

Decomposition of the velocity gradient provides a symmetric and an antisymmetric component, with the symmetric part being the strain rate tensor. This strain rate tensor is defined as $S = \frac{1}{2}(V + V^T)$. Decomposition of S yields the eigenvalues and eigenvectors of the deformation, where eigenvectors denote the principle axes of the deformation and eigenvalues the principle components of the strain rate tensor. The eigenvalues sign accounts for compressive (negative) or tensile (positive) strain. We observed very little tensile strain inside the network along the major axis. Therefore, for visualisation purposes, only the principal strain denoting compression was shown. For visual representation of control cells the principal strain field was normalised between -1 and 0. For comparing the principal strain field between genotypes, no normalisation was performed because of the reduction of the strain

values in the mutant conditions, however the colourmap scaling was fixed between genotypes.

Divergence and network turnover analysis

For quantification of divergence a central difference scheme was implemented to compute the spatial derivatives of the actin flow velocities ($\nabla \cdot V$). This method of calculating divergence was also utilised in the computation of network turnover, to determine the spatial distribution of network assembly and disassembly which was calculated using the equation below.

$$turnover = \frac{\partial I}{\partial t} + (I * (\nabla \cdot V)) + (V * (\nabla \cdot I))$$

The temporal derivative of the fluorescence intensity ($\frac{\partial I}{\partial t}$) was computed using a forward difference scheme between two consecutive frame of the time-lapse. As with the spatial gradients of flow velocity ($\nabla \cdot V$) the fluorescence intensity ($\nabla \cdot I$) was computed using a central difference scheme. As there was not much assembly information inside the lamellae of hemocytes, only the disassembly data were visually represented and normalised to the maximum value in the field. However, both assembly and disassembly was included in the quantification.

For visualisation purposes in control cells, normalised disassembly or negative divergence maps were shown normalised to the maximum value in the field, providing values between -1 and 0. For comparing the disassembly or negative divergence between genotypes, no normalisation was performed because of the reduction of these values in the mutant conditions, however the colourmap scaling was fixed between genotypes.

Linescans were used to show the contractile and destructive gradients of the flowing actin network (MATLAB *improfile* function) by drawing three random lines of 1 pixel width per frame. Lines originate from the centroid of the nucleus and extend through the lamella to the cell edge. Data points within the cell body were discarded.

Flow alignment

For determining the average alignment of actin flows, the cosine similarity between all velocity vectors and their 8 nearest neighbours was computed using $\cos\theta = v_1 \cdot v_2 / |v_1||v_2|$, and subsequently averaged to give flowfield alignment.

Extension/retraction analysis

For the extension/retraction analysis, segmented time-lapse images of freely moving hemocytes were subtracted from the subsequent frames in the time-lapse series to highlight regions of extension or retraction. The MATLAB *regionprops* function was used to filter extensions and retractions by their respective area, and to attain their centroid for the purpose of tracking these regions with respect to the position of the nucleus. Maximum extensions and retractions were defined as the regions for each frame with the largest area.

Edge velocity analysis

To evaluate edge dynamics, segmented time-lapse images of hemocytes were analysed using a Segmentation and Windowing package²⁶, calculating edge extensions and retractions at each pixel along the cell boundary. Custom scripts implemented in MATLAB were used to calculate extension speed globally, and locally within specific regions of the cell boundary. To calculate the edge velocity in the direction of cell motion, the edge was segmented within a region bounded by a 30° cone centred on the direction of motion. To calculate the edge velocity within the maximum extension, the longest uninterrupted region along the perimeter of the cell edge was segmented. To quantify the average net edge activity, positive and negative sign was assigned to velocity vectors depending on whether they were classified as extension or retractions.

Temporal cross-correlation

Temporal cross-correlation was employed to evaluate whether there was any temporal hierarchy governing the dynamics of the considered migratory parameters (i.e. cell motion, primary sink, maximum edge extension). This analysis involves the directional correlation of two vectors at all potential time lags. The temporal cross-correlation function is described as $DC = \langle v_i(t) \cdot v_j(t + \tau) \rangle$, where DC is the time

averaged cosine similarity between vector i (v_i) and parameter j (v_j) at time t and lagged time intervals ($t + \tau$).

Statistics and reproducibility

When example images are shown in figures, these represent similar results obtained from 9 independent biological samples for *wild-type* (Fig. 1-7, Extended Data Fig. 1,2,5,6), *myosin-II* (Fig. 5,6), *cofilin* (Fig. 5,6), and *ena* mutants (Extended Data Fig. 5); from 4 independent biological samples for directed migration (Fig. 3, Extended Data Fig. 4); from 5 independent biological samples for actin and myosin-II analysis (Fig. 7, Extended Data Fig. 6); from 3 independent biological samples for RPE1 cells (Extended Data Fig. 3); from 3 independent biological samples for neuronal growth cones (Extended Data Fig. 3); from 2 independent biological samples for fish keratocytes (Extended Data Fig. 3).

References

26. Machacek, M. & Danuser, G. Morphodynamic profiling of protrusion phenotypes. *Biophys J* **90**, 1439-1452 (2006).
30. Davis, J.R. *et al.* Inter-cellular forces orchestrate contact inhibition of locomotion. *Cell* **161**, 361-373 (2015).
65. Zanet, J. *et al.* Fascin promotes filopodia formation independent of its role in actin bundling. *J Cell Biol* **197**, 477-486 (2012).
66. Millard, T.H. & Martin, P. Dynamic analysis of filopodial interactions during the zippering phase of *Drosophila* dorsal closure. *Development* **135**, 621-626 (2008).
67. Franke, J.D., Montague, R.A. & Kiehart, D.P. Nonmuscle myosin II generates forces that transmit tension and drive contraction in multiple tissues during dorsal closure. *Curr Biol* **15**, 2208-2221 (2005).
68. Behrndt, M. *et al.* Forces driving epithelial spreading in zebrafish gastrulation. *Science* **338**, 257-260 (2012).
69. Dix, C.L. *et al.* The Role of Mitotic Cell-Substrate Adhesion Re-modeling in Animal Cell Division. *Dev Cell* **45**, 132-145 e133 (2018).
70. Stramer, B. *et al.* Live imaging of wound inflammation in *Drosophila* embryos reveals key roles for small GTPases during in vivo cell migration. *J Cell Biol* **168**, 567-573 (2005).

Figure 1

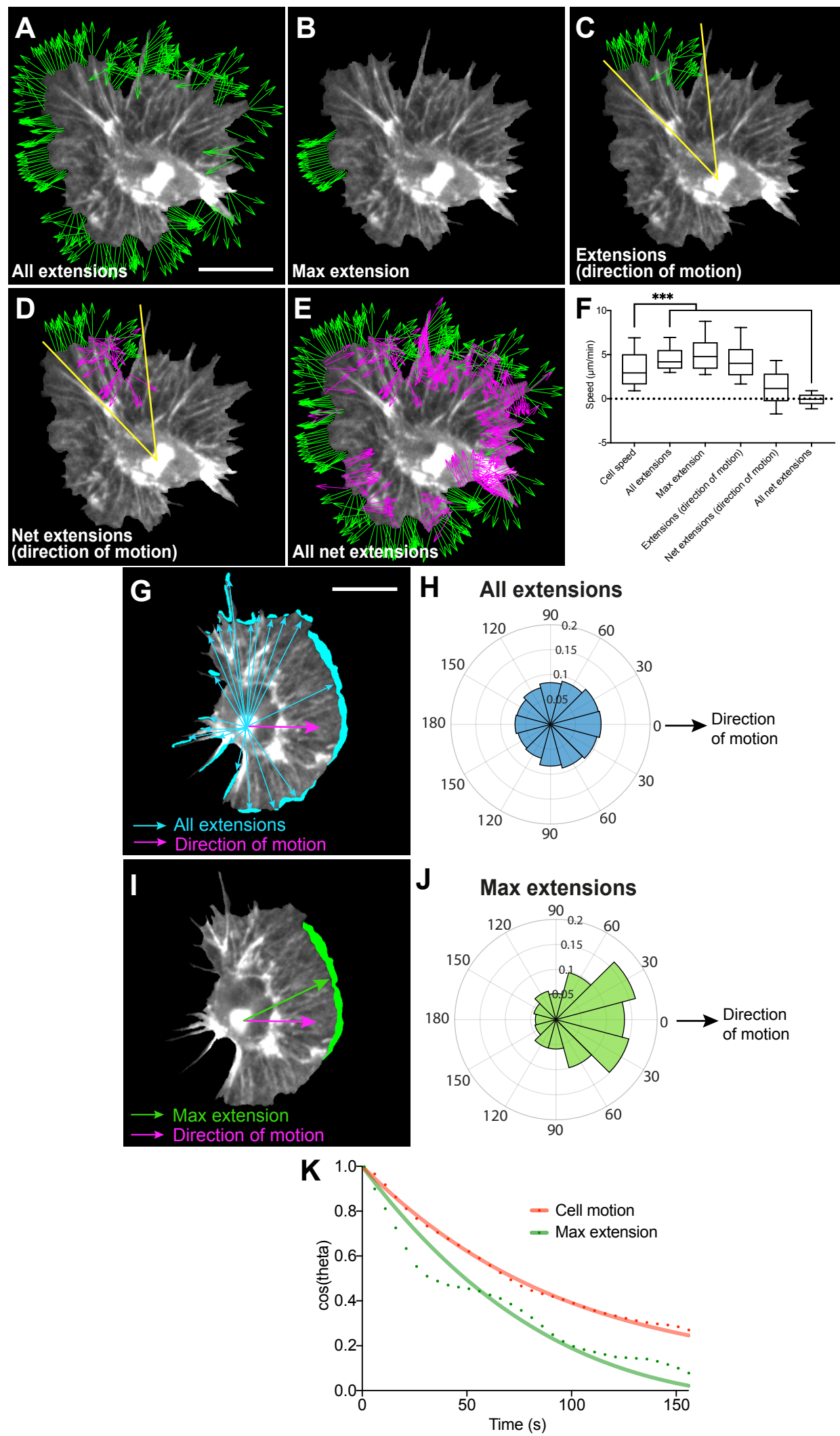


Figure 2:

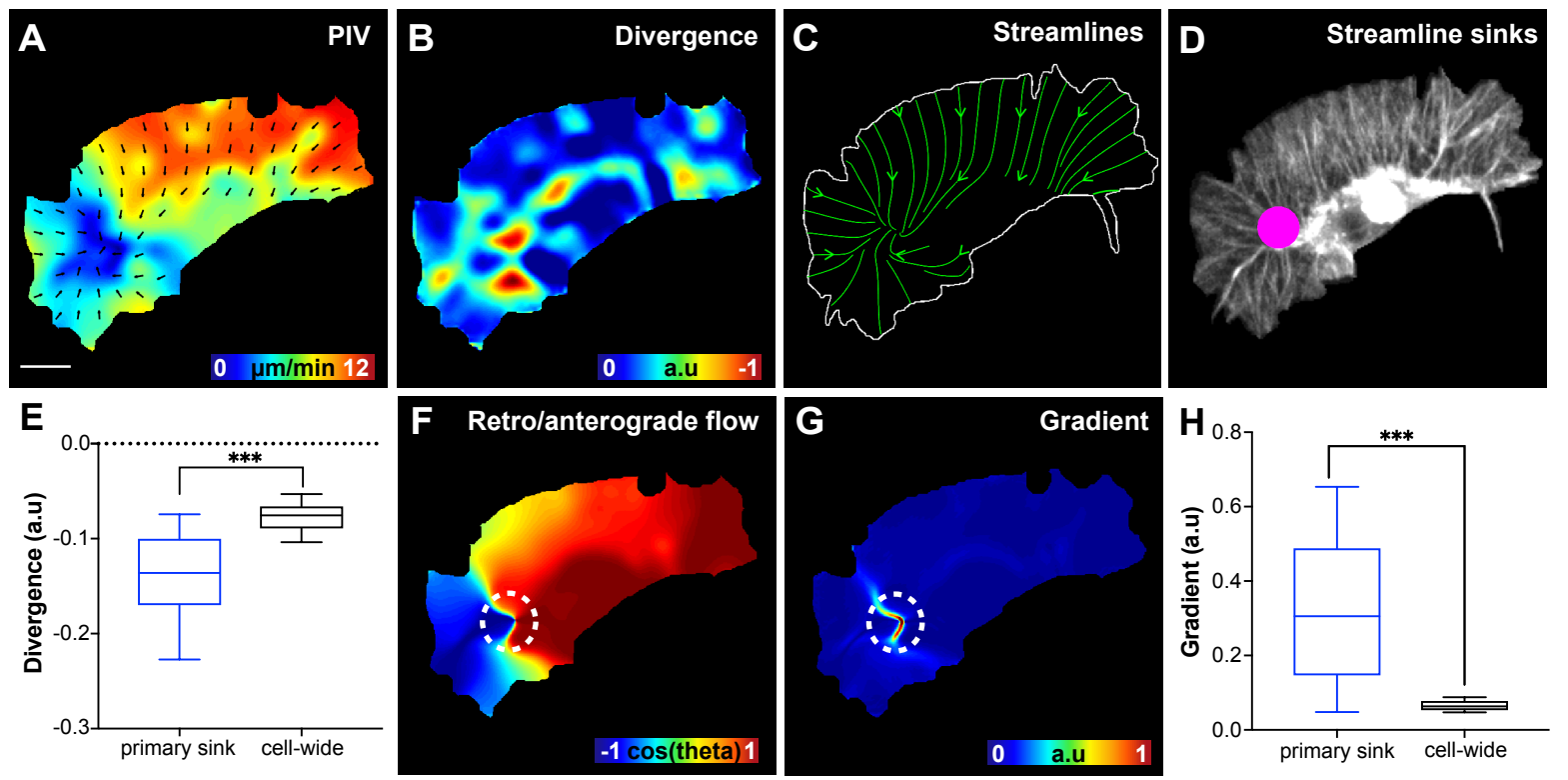


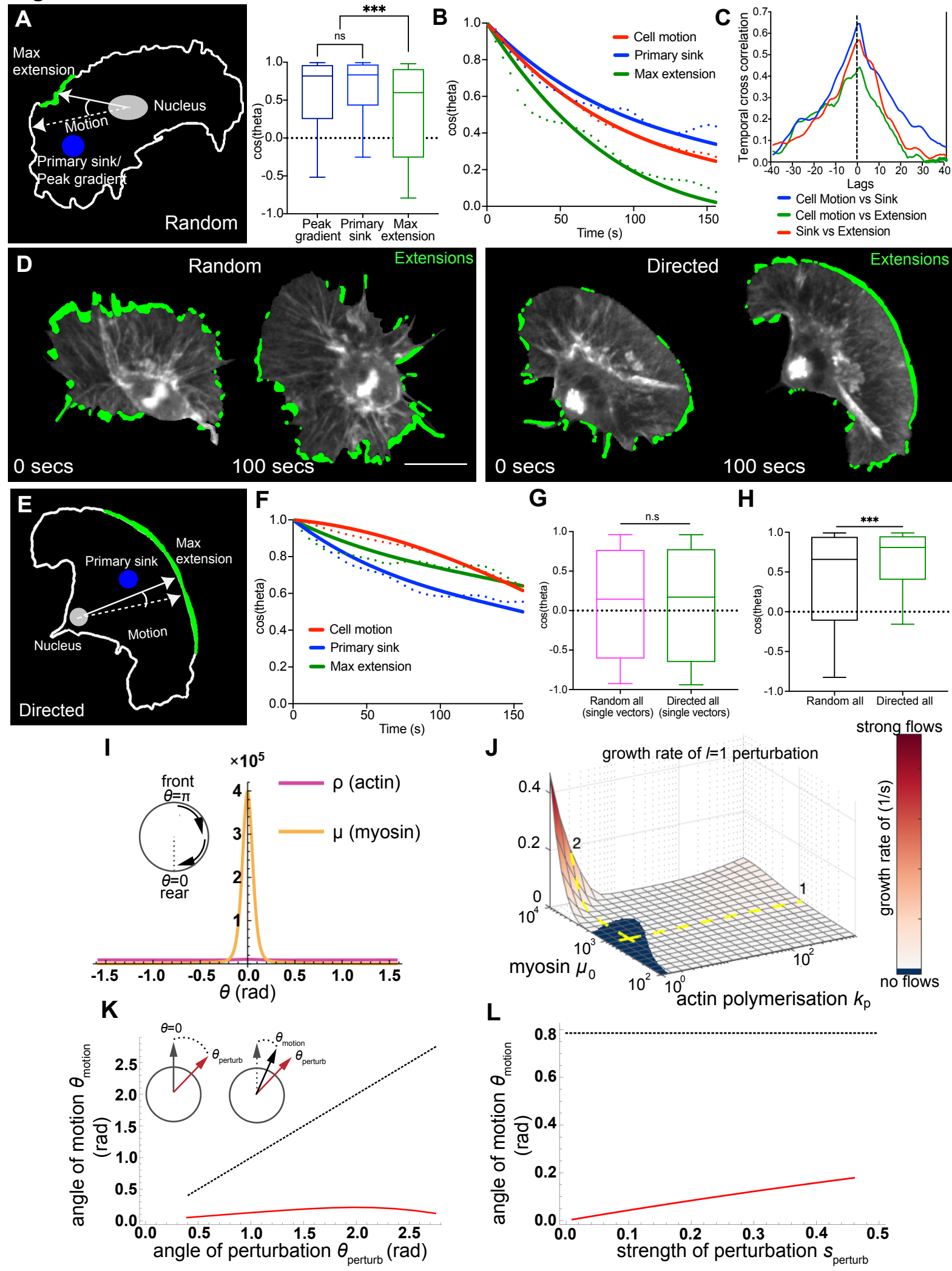
Figure 3:

Figure 4:

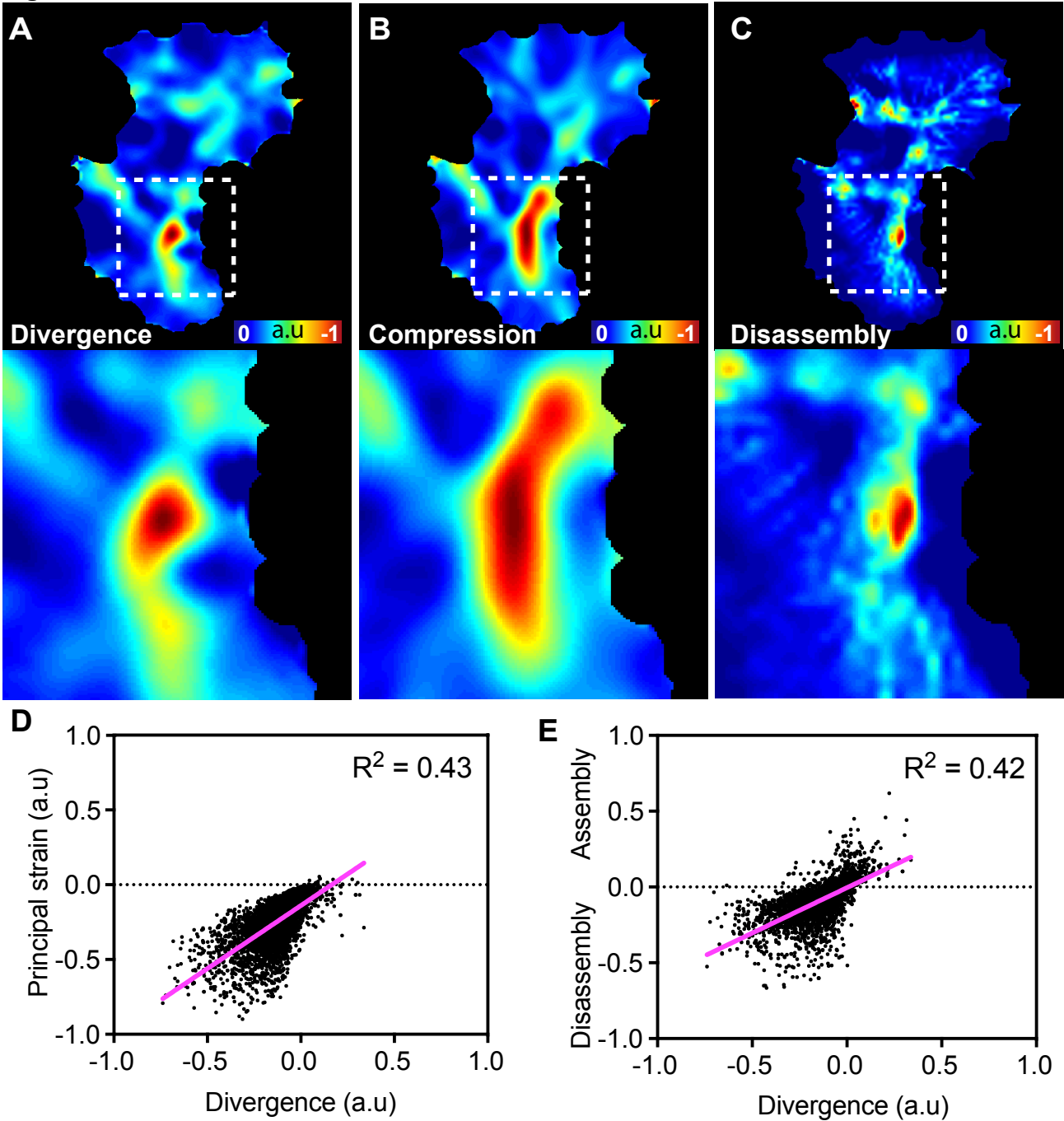


Figure 5:

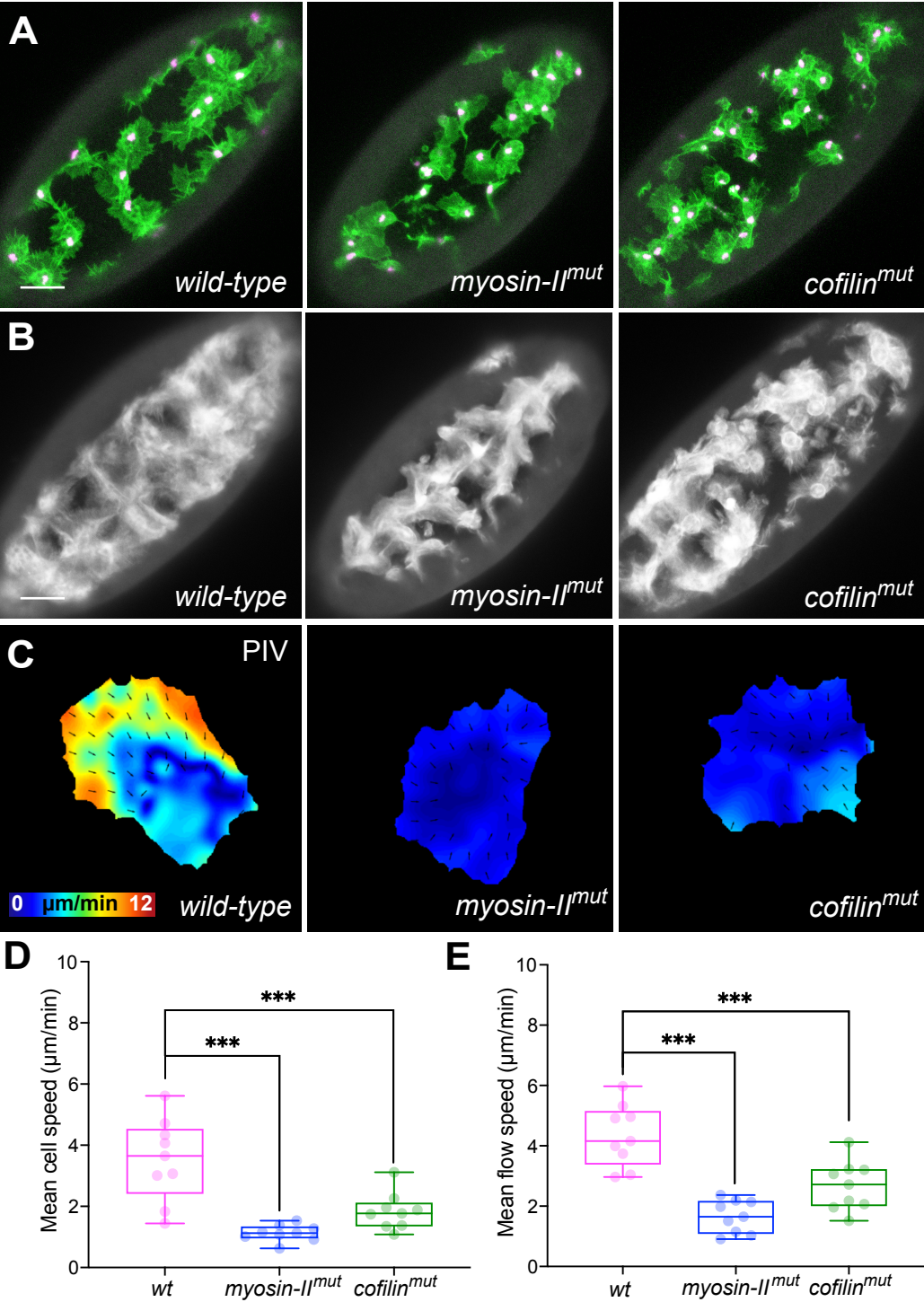


Figure 6:

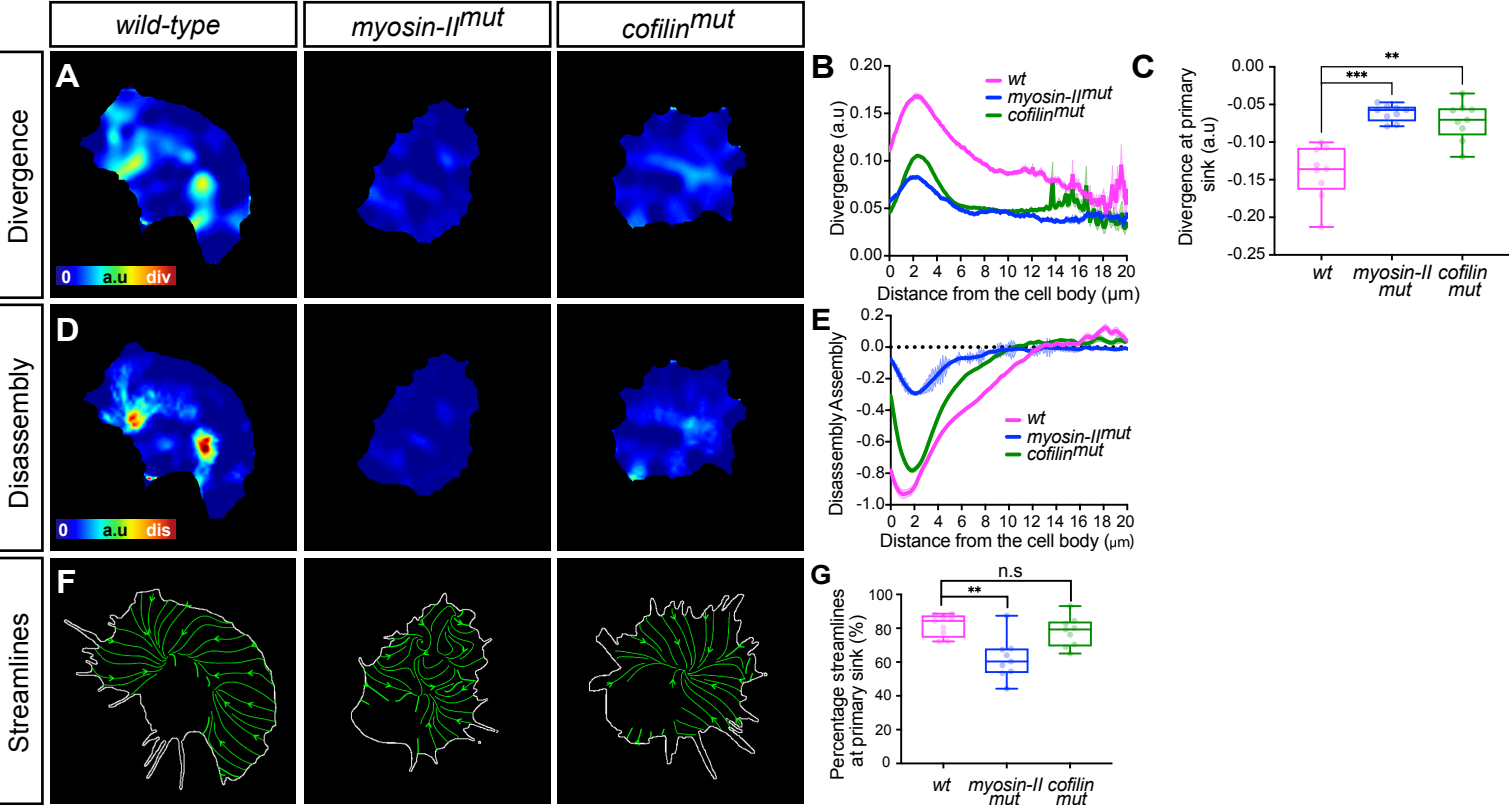


Figure 7:

